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(54) Title: C. ANTARCTICA LIPASE AND LIPASE VARIANTS

(57) Abstract

A lipase variant of a parent lipase comprising a trypsin-like catalytic triad including an active serine located in a predominantly hydrophobic, elongated binding pocket of the lipase molecule and, located in a critical position of a lipid contact zone of the lipase structure, an amino acid residue different from an aromatic amino acid residue, which amino acid residue interacts with a lipid substrate at or during hydrolysis, in which lipase variant said amino acid residue has been replaced by an aromatic amino acid residue so as to confer to the variant an increased specific activity as compared to that of the parent lipase. The parent lipase may be a C. antarctica lipase A essentially free from other substances from C. antarctica, which comprises the amino acid sequence shown in SEQ ID No. 2, or a variant of said lipase which (1) has lipase activity, (2) reacts with an antibody reactive with at least one epitope of C. antarctica lipase A having th eamino acid sequence SEQ ID No. 2, and/or (3) is encoded by a nucleotide sequence which hybridizes with an oligonucleotide probe prepared on the basis of the full or partial nucleotide sequence shown in SEQ ID No. 1 encoding the C. antarctica lipase A.

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C. antarctica lipase and lipase variants.

FIELD OF THE INVENTION

The present invention relates to novel lipase enzyme variants with improved properties, DNA constructs coding for the expression of said variants, host cells capable of expressing the variants from the DNA constructs, as well as a method of producing the variants by cultivation of said host cells. Furthermore, the present invention relates to a recombinant essentially pure <u>Candida antarctica</u> lipase and variants thereof as well as a DNA sequence encoding the said lipase or variants thereof.

BACKGROUND OF THE INVENTION

A wide variety of lipases of microbial and mammalian origin are known. The amino acid sequence of many of these lipases have 15 been elucidated and analyzed with respect to structural and functional elements important for their catalytic function, see, for instance, Winkler et al., 1990 and Schrag et al., 1991. It has been found that the lipase enzyme upon binding of a lipid substrate and activation undergoes a conformational change, which inter alia, results in an exposure of the active site to the substrate. This conformational change together with the presumed interaction between enzyme and substrate have been discussed by, inter alia, Brady et al., 1990, Brzozowski et al., 1991, Derewenda et al., 1992.

25 Based on the knowledge of the structure of a number of lipases, it has been possible to construct lipase variants having improved properties by use of recombinant DNA techniques. Thus, WO 92/05249 discloses the construction of certain lipase variants, in which the lipid contact zone has been modified so as 30 to provide the variants with different substrate specificities and/or an improved accessibility of the active site of the

- lipase to a lipid substrate. The modifications involve changing the electrostatic charge, hydrophobicity or the surface conformation of the lipid contact zone by way of amino acid substitutions.
- 5 Although the structural and functional relationship of lipases have been the subject of a number of studies as described in the above cited references, the research has mainly focused on the macroscopic characteristics of the lipases upon substrate binding and activation, whereas the identity of the amino acids 10 actually involved in the substrate binding and catalytic activity has been discussed only to a lesser extent.

SUMMARY OF THE INVENTION

By sequence alignment analysis combined with analysis of the structure and activity of a number of lipases, the present inventors have now surprisingly found that the presence of certain amino acids, especially tryptophan, in a critical position of the lipase seems to be important for optimal catalytic activity.

It is consequently an object of the present invention to modify 20 lipases which do not comprise such an amino acid residue in the critical position (which lipases in the present context are termed parent lipases) by replacing the amino acid residue located in this position with an amino acid residue which gives rise to a variant having an increased specific activity.

25 More specifically, in one aspect the present invention relates to a lipase variant of a parent lipase comprising a trypsinlike catalytic triad including an active serine located in a predominantly hydrophobic, elongated binding pocket of the lipase molecule and, located in a critical position of a lipid 30 contact zone of the lipase structure, an amino acid residue different from an aromatic amino acid residue, which interacts with a lipid substrate at or during hydrolysis, in which lipase variant said amino acid residue has been replaced by an aromatic amino acid residue so as to confer to the variant an increased specific activity as compared to that of the parent slipase.

In the present context, the term "trypsin-like" is intended to indicate that the parent lipase comprises a catalytic triad at the active site corresponding to that of trypsin, i.e. the amino acids Ser, His and one of Asp, Glu, Asn or Gln.

10 Lipases degrades triglycerides down to fatty acids, glycerol and di- and/or monoglycerides. The lipase action is depending on interfacial activation of the lipase in the presence of substrate surfaces. On activation lipases change their conformation in such a manner that their surface hydrophobicity in an 15 area around the active site is increased. The interfacial activation of lipases is e.g. discussed by Tilbeurgh et al. (1993).

All lipases studied until now have been found to comprise at least one surface loop structure (also termed a lid or a flap) 20 which covers the active serine when the lipase is in inactive form (an example of such a lipase is described by Brady et al., 1990). When the lipase is activated, the loop structure is shifted to expose the active site residues, creating a surface surrounding the active site Ser, which has an increased surface 25 hydrophobicity and which interacts with the lipid substrate at or during hydrolysis. For the present purpose, this surface is termed the "lipid contact zone", intended to include amino acid residues located within or forming part of this surface, optionally in the form of loop structures. These residues may 30 participate in lipase interaction with the substrate at or during hydrolysis where the lipase hydrolyses triglycerides from the lipid phase when activated by contact with the lipid surface.

The lipid contact zone contains a binding area (a so-called binding pocket) for the lipid substrate which is the part of the lipid contact zone to which the lipid substrate binds before hydrolysis. This binding area again contains a so-called hydrolysis pocket, which is situated around the active site Ser, and in which the hydrolysis of the lipid substrate is believed to take place. In all known lipases to day the lipid contact zone is easily recognized, e.g. from a three-dimensional structure of the lipase created by suitable computer programs. The conformation of an inactive and activated lipase, respectively, is shown in Fig. 1 which is further discussed below.

In the present context, the "critical position" of the lipase molecule is the position in the lipid contact zone of the lipase molecule, which is occupied by an amino acid residue which interacts with the lipid substrate and which is different from an aromatic amino acid residue.

In another aspect the present invention relates an <u>C. antarctica</u> lipase A which is essentially free from other <u>C. antarctica</u> substances and which comprises the amino acid sequence identified in SEQ ID No. 2 or a variant thereof which

- 1) has lipase activity,
- 2) reacts with an antibody reactive with at least one epitope of the <u>C. antarctica</u> lipase having the amino acid sequence 25 shown in SEQ ID No. 2, and/or
 - 3) is encoded by a nucleotide sequence which hybridizes with an oligonucleotide probe prepared on the basis of the full or partial nucleotide sequence shown in SEQ ID No. 1 encoding the <u>C. antarctica</u> lipase A.
- 30 The <u>C. antarctica</u> lipase A of the invention has a number of desirable properties including a high thermostability and

activity at acidic pH and may advantageously be produced by use of recombinant DNA techniques, e.g. using the procedures described below. Thus, the lipase A of the invention may be obtained in a higher purity and a higher amount than the <u>C. antarctica</u> lipase A purified from wild type <u>C. antarctica</u> which is described in WO 88/02775.

Furthermore, the present invention relates to a DNA sequence encoding the <u>C. antarctica</u> lipase A having the amino acid sequence identified in SEQ ID No. 2 or a modification of said 10 DNA sequence encoding a variant of the <u>C. antarctica</u> lipase A as defined above.

In the present context "C. antarctica lipase A" is used interchangeably with "lipase A" and the variant of the C. antarctica lipase A is termed "lipase A variant".

15 The present invention also relates to a DNA construct comprising a DNA sequence encoding a lipase variant as indicated above or a DNA sequence encoding the <u>C. antarctica</u> lipase A, a recombinant expression vector carrying said DNA construct, a cell transformed with the DNA construct or the expression vector, as well as a method of producing a lipase variant of the invention by culturing said cell under conditions conducive to the production of the lipase variant, after which the lipase variant is recovered from the culture.

It will be understood that lipase variants of the present invention having an increased specific activity as compared to their parent lipases may be used for the same purposes as their parent lipases, advantageously in a lower amount due to their higher specific activity.

Accordingly, the present invention relates to the use of a 30 lipase variant of the invention as a detergent enzyme; as a digestive enzyme; in ester hydrolysis, ester synthesis or interesterification; or the use of the lipase variant to avoid pitch

trouble arising, e.g., in processes for preparing mechanical pulp and in paper-making processes using mechanical pulp.

DETAILED DISCLOSURE OF THE INVENTION

As indicated above, the present inventors have found that the presence of certain aromatic amino acids, especially tryptophan, located in the lipid contact zone of the lipase molecule is important for optimal catalytic activity.

The importance of the presence of an aromatic amino acid residue and in particular a tryptophan residue was found in connection with a study of mutants of a <a href="https://www.humicola.com/humicola.

Without being limited to any theory it is presently believed that the amino acid residue present in the critical position, e.g. on top of or in the proximity of the active serine, may be involved in a) stabilization of the tetrahedral intermediate formed from the lipase and the substrate during the activation of the lipase, and b) in the activation of the replacement of the lid region covering the active serine in the inactive enzyme. When tryptophan is present in this position, it is contemplated that optimal performance with respect to a) as well as b) above is obtained. Thus, it is believed that tryptophan gives rise to the formation of the most stable tetrahedral intermediate (which means a lowering of the activation energy

needed for the catalysis to take place), and further improves the performance of the enzyme with respect to the activation of the lid opening which is essential for any catalysis to take place.

- 5 In connection with a) above it has been observed that the best acting lipase variants contain an unsaturated ring system in the side-chain. The far the biggest unsaturated system is tryptophan, then tyrosine, phenylalanine and histidine. These sidechains have a pi-electron system ("the unsaturation") that could be important for the proton transfer in the catalysis resulting in a lower activation energy for creating the tetrahedral intermediate where proton transfer has taken place from active site histidine to serine to the oxyanion hole created after lid activation and opening.
- 15 From the above theoretical explanation it will be understood that the optimal amino acid to be present in the critical position, e.g. on top of or in the proximity of the active serine, is tryptophan. However, when the parent lipase is one which does not contain any aromatic amino acid residue or any 20 amino acid residue with an unsaturated ring system in the sidechain in this position, such amino acids may advantageously be substituted into this position.

Thus, when the parent lipase, in the critical position, has an amino acid residue which does not comprise an unsaturated ring system in the side-chain, an amino acid residue having such an unsaturated ring-system, e.g. an aromatic amino acid (tryptophan, tyrosine, phenylalanine or histidine) may be substituted into the critical position. When the amino acid residue in the critical position of the parent lipase is histidine, it may advantageously be replaced by phenylalanine, tyrosine and most preferably tryptophan, when the amino acid residue is tyrosine, it may advantageously be replaced by phenylalanine and most preferably tryptophan, and when the

pmino acid residue is phenylalanine it may advantageously be replaced by tryptophan.

While the critical position in some lipases is contemplated to be any position within the lipid contact zone, the critical sposition will normally be located in the binding pocket of the lipase molecule, and preferably in the hydrolysis pocket thereof. For most lipases it is believed that the critical amino acid residue is positioned on top of or in the proximity of the active site.

The amino acid residue occupying this position may be identified in any lipase by 1) sequence alignment studies in which the amino acid sequence of the lipase in question is aligned with the amino acid sequence of other lipases, in which the amino acid residue positioned on top of or in the proximity of 15 the active serine has been identified, so as to identify the presumed position of said amino acid residue, and/or 2) an analysis of the three-dimensional structure of the lipase in question using standard display programmes such as INSIGHT (Biosym Technologies Inc., San Diego, USA), so as identify the 20 amino acid sequence on top of or in the proximity of the active serine.

More specifically, on the basis of a computer program such as INSIGHT displaying lipase coordinates in accordance with well-known technology, it is simple to point out which part of the 25 lipase which contains the lipid contact zone. 1/ if the structure of the lipase is in a non-activated form, the lipid contact zone is identified by the direction of sidechains of the active site Ser. 2/ if the structure is in the activated form one may additionally base the identification on a colour-ing of all hydrophobic residues in a colour different from the other residues. By this procedure in which a cpk model of the structure is created, the hydrophobic surface specific for the lipid contact zone may be identified. The active site Ser is located within this more hydrophobic part of the molecule.

In some lipases the critical amino acid residue is located in the surface loop structure covering the active site, or in one or more of the surface loop structures found to form part of the surface of the lipid contact zone, such as of the binding 5 pocket or hydrolysis pocket.

Although the critical position is normally considered to be constituted of only one amino acid residue it may be advantageous to replace two or more residues, preferably with a tryptophan residue as explained above, in order to obtain a 10 further increased specific acitivity.

It is contemplated that it is possible to increase the specific activity of parent lipases which do not have a tryptophan residue in the critical position at least 2 times, such as at least 3 and preferably at least 4 or even 5, 6 or 7 times by 15 modifications as diclosed herein.

It is contemplated that lipase variants as defined herein having an increased substrate specificity may be prepared on the basis of parent lipases of various origins. Thus, the parent lipase may be a microbial lipase or a mammalian lipase.

20 When the parent lipase is a microbial lipase, it may be selected from yeast, e.g. <u>Candida</u>, lipases, bacterial, e.g. <u>Pseudomonas</u>, lipases or fungal, e.g. <u>Humicola</u> or <u>Rhizomucor</u> lipases.

One preferred lipase variant is one, in which the parent lipase is derived from a strain of <u>Candida antarctica</u>, in particular one in which the parent lipase is lipase A of <u>C. antarctica</u>, preferably the one which has the amino acid sequence shown in SEQ ID No. 2 or a lipase A variant thereof as defined herein. The lipase variant of this <u>C. antarctica</u> lipase A preferably has the amino acid sequence shown in SEQ ID No. 2 in which the phenylalanine 139 of the parent lipase has been replaced by a tryptophan residue. The construction of this variant and the

analysis of the properties thereof is discussed in Example 3, 5 and 6.

A lipase variant of the invention may, as mentioned above, be prepared on the basis of a parent lipase derived from a strain of a <u>Pseudomonas</u> species, e.g. <u>Ps. fragi</u>. An example of a suitable <u>Ps. fragi</u> lipase which has an amino acid residue different from tryptophan positioned on top of or in the proximity of the active serine, is the one described by Aoyama et al., 1988. A lipase variant according to the present invention may be constructed by replacing the phenylalanine residue 29 in the amino acid sequence of said lipase shown in SEQ ID No. 3 by a tryptophan residue.

An example of a fungal lipase suitable as a parent lipase for the construction of a lipase variant of the invention is one derived from Rhizopus, especially from R. delemar or R. niveus, the amino acid sequence of which latter is disclosed in, e.g., JP 64-80290. In order to construct a lipase variant according to the present invention from this parent lipase, the alanine residue at position 117 is to be replaced with an aromatic amino acid residue such as tryptophan. The sequence alignment of the R. niveus lipase sequence (SEQ ID No. 5) and an Rhizomucor miehei lipase sequence (containing a tryptophan residue) (SEQ ID No. 4) is illustrated below. From this alignment the critical position of the R. niveus lipase may be determined.

25 SEQUENCE		10		20		30		40		50	60	Res#
						1						L
mucor						-sidg	GIRA	ATSQEI	NELIT	YYTTLS	SANSYCRU	V 32
niveus	DONLVG	MILDL	PSDAI	PISLS	SSINS	SASDGG	KVVA	ATTAQI	QEFT	KYAGI	ATAYCRS	V 60
		70		80		90		100		110	10	0 Res#
SEQUENCE		70		80		90		100		110		·
30												i.
mucor	IPGAIW	DCIHCI	ATE-I	DIKIIK	IWSI	LIYDIN	IAMVA	RCDSEK	TYI	VFRGS	SSIRNWIA	D 91
niveus	VPGNKW	DCVQQQ	KWP	OGKIII	TFTS	LLSDIN	IGYVI	RSDKQK	IXIE	VFRGI	NSFRSATI	D 120

SEQUENCE	130	140	150	160	170	180 Res#
mucor niveus	LITYPVSYPPVSIVFNFSDYKPVK					
5 SEQUENCE	190	200	210	220	230 	240 Res#
mucor niveus	LCALGLYOREEG LAGMOLYOREPR					
SEQUENCE	250	260	270	280	290	300 Res#
mucor niveus	AFGFIHAGEEYV SFGFIHPGVESV					

The present inventors have surprisingly found that non-pancreatic lipases such as gastric, lingual, or hepatic lipases 15 have the common feature that the amino acid residue which has been identified to be the one located in the critical position of the lipase molecule, normally on top of or in the proximity of the active serine, is different from tryptophan. This is in contrast to pancreatic lipases which generally have been found 20 to have a tryptophan residue in this position. Thus, in the present context, non-pancreatic mammalian lipases may advantageously be used as "parent lipases" for the construction of lipase variants of the invention.

Accordingly, lipase variants as disclosed herein which is of 25 mammalian origin is advantageously prepared from a parent lipase of non-pancreatic, such as gastric, lingual or hepatic origin. Such mammalian lipases may be derived from humans, rats, mice, pigs, dogs or other mammals. Specific examples of such mammalian lipases includes a rat lingual lipase having the sequence identified as A23045 (Docherty et al., 1985), a rat hepatic lipase having the sequence identified as A27442

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(Komaromy and Schotz, 1987), a human hepatic lipase having the sequence identified as A33553 (Datta et al., 1988), a human gastric lipase having the sequence identified as S07145 (Bodmer et al., 1987), and a Bio Salt Activated Lipase (BSAL) having the sequence identified as A37916 (Baba et al., 1991) all of which were analysed with respect to the critical position in the sequence alignment analysis illustrated below. The pancreatic lipases included in this sequence alignment study were a murine pancreatic lipase, A34671 (Grusby et al., 1990), a porcine pancreatic lipase, A00732 (Caro et al., 1981), a human pancreatic lipase, A34494 (Lowe et al., 1989), and a canine pancreatic lipase having the sequence B24392 (Mickel et al., 1989). The amino acid sequences of each of the lipases mentioned have the accession numbers listed above and are available from publically available databases.

A37916 TYGDEDCLYL NIWVPQGRK. ..QVSRDLPV MIWIYGGAFL MGSGHGANFL A23045 EVVTEDGYIL GVYRIPHGKN NSENIGKRPV VYLQHGLIAS AT..NWIANL 507145 EVVTEDGYIL EVNRIPYGKK NSCNIGORPV VFLOHGLIAS AT..NWISNL B24392 TNKNPNNFQT LLPSDPSTIE ASNFQIDKKT RFTIHGFINK GE.ENWLLDM 20 TNENPNNFQE VA.ADSSSIS GSNFKINRKT RFIIHGFIDK GE.ENWLANV A34494 A34671 TNENPNNYQI ISATDPATIN ASNFQLDRKT RFTIHGFIDK GE.EGWLLDM A00732 TNONONNYOE LV.ADPSTIT NSNFRMDRKT RFIIHGFIDK GE.EDWLSNI A33553 GEINQ..GOQ IRINHPDIIQ EOGFNSSLPL VMIIHGWSVD GVLENWIWQM A27442 KDESDRLGOQ IRPOHPETLQ EOGFNSSHPL VMIIHGWSVD GLLETWIWKI 25 130 90 A37916 NNYLYDGEEI ATRCNVIVVI FNYRVGPLGF LSTGDANLPG NYGLRDQHMA A23045 PNNSLAFMLA DAGYDVWLGN SRGNTWSRKN VYYSPDSVEF WAFSFDEMAK 507145 PNNSLAFTLA DAGYDVWLGN SRGNTWARRN LYYSPDSVEF WAFSFDEMAK B24392 CKNMFKVEE.VN CICVDWKKGS QTSYTQAANN VRVVGAQVAQ 30 A34494 CKNLFKVES.VN CICVDWKGGS RIGYTQASQN IRIVGAEVAY A34671 CKKMFQVEK.VN CICVDWKRGS RIEYTQASYN TRVVGAEIAF A00732 CKNLFKVES.VN CICVDWKGGS RIGYTQASQN IRIVGAEVAY A33553 VAALKSOPAO P......VN VGLVDWITLA HDHYTIAVRN TRLVGKEVAA A27442 VGALKSRQSQ P......VN VGLVDWISLA YQHYAIAVRN TRVVGQEVAA 35

è		131	į		1	175
	A37916	IAWVKRNI.A	AFGGDPNNIT	LFGESAGGAS	VSLQTLSPYN	KGLIRRA
	A23045	YDLPATINFI	VQKTGQEKTH	YVGHSQGTTI	GFIAFSINPT	LAKKIKIT
	S07145	YDLPATIDFI	VKKTGQKQLH	YVGHSQGITI	GFIAFSINPS	LAKRIKIF
5	B24392	MLSMLSA	NYSYSPSQVQ	LIGHSLGAHV	AGEAGSRIPG	LGRITGL
	A34494	FVEFLQS	AFGYSPSNVH	VIGHSLGAHA	AGEAGRRING	TIGRITGL
	A34671	LVQVLST	EMGYSPENVH	LIGHSLGSHV	AGEAGRRLEG	HVGRITGL
	A00732	FVEVLKS	SLGYSPSNVH	VIGHSLGSHA	AGEAGRRING	TIERITGL
	A33553	LIRWLEE	SVQLSRSHVH	LIGYSLGAHV	SGFAGSSIGG	THKIGRITGL
10	A27442	LLLWLEE	SMKFSRSKVH	LIGYSLGAHV	SGFAGSSMGG	KKKIGRITGL
		•		L	I	
		176				220
	A37916	ISQSGVALSP	WVIQKN	PLFWAKKV	AEKVGCPVGD	AARMAQCLKV
15	A23045	YALAPVATVK	YTOSPLKKIS	FIPIFLFKIM	FGKKMFLPHT	YFDDFLGTEV
	S07145	YALAPVATVK	YTKSLINKLR	FVPQSLFKFI	FGDKIFYPHN	FFDQFLATEV
	B24392	DPVEASFQGT	PEEVRLD	PIDADFVD	VIHIDAAPLI	PFLGFGTSQQ
	A34494	DPAEPCFQGT	PELVRLD	PSDAKFVD	VIHIDGAPIV	PNLGFGMSQV
	A34671	DPAEPCFQGL	PEEVRLD	PSDAMFVD	VIHIDSAPII	PYLGFGMSQK
20	A00732	DPAEPCFQGT	PELVRLD	PSDAKFVD	VIHIDAAPII	PNLGFGMSQT
	A33553	DAAGPLFEGS	APSNRLS	PDDASFVD	AIHIFTREHM	GLSVGIK.QP
	A27442	DPAGPMFEGT	SPNERLS	PDDANFVD	AIHIFIREHM	GLSVGIK.QP
				Z = Flap	region	
25		221				27 0
	A37916	TOPRALITLAY	KVPLAGLEYP	MLHYVGFVPV	IDGDFIPADP	INLYANAADI
	A23045	CSREVLDLLC	SNILFIFCGF	DKKNLNVSRF	DVYLGHNPAG	TSVQDFLHWA
	S07145	CSREMINLLC	SNALFIICGF	DSKNFNTSRL	DVYLSHNPAG	TSVQNMFHWI
	B24392	MGHLDFFPNG	GEEMPGCKKN	ALSQIVNLDG	IWEGIRDFVA	CNHLRSYKYY
30	A34494	VGHLDFFPNG	GVEMPGCKKN	TLSQIVDIDG	IWEGTRDFAA	CNHLRSYKYY
	A34671	VGHLDFFPNG	GKEIPGCOKN	ILSTIVDING	IWEGIRNFAA	CNHLRSYKYY
	A00732	VGHLDFFPNG	CKOMPGCOKN	ILSQIVDIDG	IWEGTRDFVA	CNHLRSYKYY
	A33553	ICHYDFYPNG	GSFQPGCHFL	ELYRHIAQHG	FNAITQIIK.	CSHERSVHLF
	A27442	IAHYDFYPNG	GSFQPGCHFL	ELYKHIAEHG	LNATTOTIK.	CAHERSVHLF

As mentioned above the present invention also relates to a <u>C. antarctica</u> lipase A essentially free from other <u>C. antarctica</u> substances, which has the amino acid sequence shown in SEQ ID No. 2 or a variant therof which

- 5 1) has lipase activity,
 - 2) reacts with an antibody reactive with at least one epitope of <u>C. antarctica</u> lipase A having the amino acid sequence shown in SEQ ID No. 2, and/or
- 3) is encoded by a nucleotide sequence which hybridizes with an 10 oligonucleotide probe prepared on the basis of the full or partial nucleotide sequence shown in SEQ ID No. 1 encoding the C. antarctica lipase A.

In the present context, the term "variant" is intended to indicate a lipase A variant which is derived from the C.

15 antarctica lipase A having the amino acid sequence shown in SEQ ID No. 2, or a naturally occurring variant. Typically, the variant differ from the native lipase A by one or more amino acid residues, which may have been added or deleted from either or both of the N-terminal or C-terminal end of the lipase,

20 inserted or deleted at one or more sites within the amino acid sequence of the lipase or substituted with one or more amino acid residues within, or at either or both ends of the amino acid sequence of the lipase.

Furthermore, the variant of the invention has one or more of 25 the characterizing properties 1)-3) mentioned above. Property 1), i.e. the "lipase activity" of the variant may be determined using any known lipase assay, e.g. the Standard LU assay described in the Methods section below.

Property 2), i.e. the reactivity of the variant of the inven-30 tion with an antibody raised against or reactive with at least one epitope of the <u>C. antarctica</u> lipase A having the amino acid sequence shown in SEQ ID No. 2 below may be determined by polyclonal antibodies produced in a known manner, for instance by immunization of a rabbit with the <u>C. antarctica</u> lipase A of the invention. The antibody reactivity may be determined using assays known in the art, examples of which are Western Blotting or radial immunodiffusion assay.

Property 3) above, involving hybridization, may be performed using an oligonucleotide probe prepared on the basis of the full or partial cDNA sequence encoding the <u>C. antarctica</u> lipase 10 A, the amino acid sequence of which is identified in SEQ ID No. 2, as a hybridization probe in a hybridization experiment carried out under standard hybridization conditions. For instance, such conditions are hybridization under specified conditions, e.g. involving presoaking in 5xSSC and prehy15 bridizing for 1h at ~40°C in a solution of 20% formamide, 5xDenhardt's solution, 50mM sodium phosphate, pH 6.8, and 50µg of denatured sonicated calf thymus DNA, followed by hybridization in the same solution supplemented with 100µM ATP for 18h at ~40°C, or other methods described by e.g. Sambrook et al., 20 1989.

The nucleotide sequence on the basis of which the oligonucleotide probe is prepared is conveniently the DNA sequence shown in SEQ ID No. 1.

As stated above in a further aspect the present invention 25 relates to a DNA sequence encoding <u>C. antarctica</u> lipase A having the amino acid sequence shown in SEQ ID No. 2 or a modification of said DNA sequence which encodes a variant of <u>C. antarctica</u> lipase A which

- 1) has lipase activity,
- 30 2) reacts with an antibody reactive with at least one epitope of the <u>C. antarctica</u> lipase A having the amino acid sequence shown in SEQ ID No. 2, and/or

- (3) is encoded by a nucleotide sequence which hybridizes with an oligonucleotide probe prepared on the basis of the full or partial nucleotide sequence shown in SEQ ID No. 1 encoding the C. antarctica lipase A.
- 5 Examples of suitable modifications of the DNA sequence are nucleotide substitutions which do not give rise to another amino acid sequence of the encoded enzyme, but which may correspond to the codon usage of the host organism into which the DNA sequence is introduced or nucleotide substitutions which do give rise to a different amino acid sequence, without, however, impairing the above stated properties of the enzyme. Other examples of possible modifications are insertion of one or more nucleotides into the sequence, addition of one or more nucleotides at either end of the sequence and deletion of one or more nucleotides at either end of or within the sequence.

Methods of preparing lipase variants of the invention

Several methods for introducing mutations into genes are known in the art. After a brief discussion of cloning lipase-encoding DNA sequences, methods for generating mutations at specific sites within the lipase-encoding sequence will be discussed.

Cloning a DNA sequence encoding a lipase

The DNA sequence encoding a parent lipase or the <u>C. antarctica</u> lipase A as defined herein may be isolated from any cell or microorganism producing the lipase in question by various 25 methods, well known in the art. First a genomic DNA and/or cDNA library should be constructed using chromosomal DNA or messenger RNA from the organism that produces the lipase to be studied. Then, if the amino acid sequence of the lipase is known, homologous, labelled oligonucleotide probes may be 30 synthesized and used to identify lipase-encoding clones from a genomic library of bacterial DNA, or from a fungal cDNA library. Alternatively, a labelled oligonucleotide probe containing sequences homologous to lipase from another strain of bacteria or fungus could be used as a probe to identify

lipase-encoding clones, using hybridization and washing conditions of lower stringency.

Yet another method for identifying lipase-producing clones would involve inserting fragments of genomic DNA into an sexpression vector, such as a plasmid, transforming lipase-negative bacteria with the resulting genomic DNA library, and then plating the transformed bacteria onto agar containing a substrate for lipase. Those bacteria containing lipase-bearing plasmid will produce colonies surrounded by a halo of clear agar, due to digestion of the substrate by secreted lipase.

Alternatively, the DNA sequence encoding the enzyme may be prepared synthetically by established standard methods, e.g. the phosphoamidite method described by S.L. Beaucage and M.H. Caruthers (1981) or the method described by Matthes et al. 15 (1984). According to the phosphoamidite method, oligonucleotides are synthesized, e.g. in an automatic DNA synthesizer, purified, annealed, ligated and cloned in appropriate vectors.

Finally, the DNA sequence may be of mixed genomic and synthetic, mixed synthetic and cDNA or mixed genomic and cDNA origin prepared by ligating fragments of synthetic, genomic or cDNA origin (as appropriate), the fragments corresponding to various parts of the entire DNA sequence, in accordance with standard techniques. The DNA sequence may also be prepared by polymerase chain reaction (PCR) using specific primers, for instance as described in US 4,683,202 or R.K. Saiki et al. (1988).

Site-directed mutagenesis of the lipase-encoding sequence
Once a lipase-encoding DNA sequence has been isolated, and desirable sites for mutation identified, mutations may be introduced using synthetic oligonucleotides. These oligonucleotides
ontain nucleotide sequences flanking the desired mutation sites; mutant nucleotides are inserted during oligonucleotide synthesis. In a specific method, a single-stranded gap of DNA, bridging the lipase-encoding sequence, is created in a vector

carrying the lipase gene. Then the synthetic nucleotide, bearing the desired mutation, is annealed to a homologous portion of the single-stranded DNA. The remaining gap is then filled in with DNA polymerase I (Klenow fragment) and the construct is ligated using T4 ligase. A specific example of this method is described in Morinaga et al. (1984). U.S. Patent number 4,760,025 discloses the introduction of oligonucleotides encoding multiple mutations by performing minor alterations of the cassette, however, an even greater variety of mutations can be introduced at any one time by the Morinaga method, because a multitude of oligonucleotides, of various lengths, can be introduced.

Another method of introducing mutations into lipase-encoding sequences is described in Nelson and Long (1989). It involves the 3-step generation of a PCR fragment containing the desired mutation introduced by using a chemically synthesized DNA strand as one of the primers in the PCR reactions. From the PCR-generated fragment, a DNA fragment carrying the mutation may be isolated by cleavage with restriction endonucleases and 20 reinserted into an expression plasmid.

Expression of lipase variants

According to the invention, a <u>C. antarctica</u> lipase A-coding sequence or a mutated lipase-coding sequence produced by methods described above or any alternative methods known in the 25 art, can be expressed, in enzyme form, using an expression vector which typically includes control sequences encoding a promoter, operator, ribosome binding site, translation initiation signal, and, optionally, a repressor gene or various activator genes. To permit the secretion of the expressed protein, nucleotides encoding a "signal sequence" may be inserted prior to the lipase-coding sequence. For expression under the direction of control sequences, a target gene to be treated according to the invention is operably linked to the control sequences in the proper reading frame. Promoter 35 sequences that can be incorporated into plasmid vectors, and

which can support the transcription of the mutant lipase gene, include but are not limited to the prokaryotic \$\beta\$-lactamase promoter (Villa-Kamaroff et al. (1978) and the tac promoter (DeBoer, et al., 1983). Further references can also be found in "Useful proteins from recombinant bacteria" (1980).

According to one embodiment a strain of <u>Bacillus</u>, e.g. <u>B. subtilis</u>, <u>B. licheniformis</u> or <u>B. lentus</u>, or a strain of <u>E. coli</u> is transformed by an expression vector carrying the lipase A or the mutated DNA. If expression is to take place in a secreting microorganism such as <u>B. subtilis</u> a signal sequence may follow the translation initiation signal and precede the DNA sequence of interest. The signal sequence acts to transport the expression product to the cell wall where it is cleaved from the product upon secretion. The term "control sequences" as defined above is intended to include a signal sequence, when is present.

The lipase or lipase variants of the invention may further be produced by using a yeast cell has a host cell. Examples of suitable yeast host cells include a strain of <u>Saccharomyces</u>, 20 such as <u>S. cerevisiae</u>, or a strain of <u>Hansenula</u>, e.g. <u>H. polymorpha</u> or <u>Pichia</u>, e.g. <u>P. pastoris</u>.

In a currently preferred method of producing lipase A or lipase variants of the invention, a filamentous fungus is used as the host organism. The filamentous fungus host organism may conveniently be one which has previously been used as a host for producing recombinant proteins, e.g. a strain of Aspergillus sp., such as A. niger, A. nidulans or A. oryzae. The use of A. oryzae in the production of recombinant proteins is extensively described in, e.g. EP 238 023.

30 For expression of lipase variants in <u>Aspergillus</u>, the DNA sequence coding for the lipase A or the lipase variant is preceded by a promoter. The promoter may be any DNA sequence exhibiting a strong transcriptional activity in <u>Aspergillus</u> and

may be derived from a gene encoding an extracellular or intracellular protein such as an amylase, a glucoamylase, a protease, a lipase, a cellulase or a glycolytic enzyme.

Examples of suitable promoters are those derived from the gene sencoding <u>A. oryzae</u> TAKA amylase, <u>Rhizomucor miehei</u> aspartic proteinase, <u>A. niger</u> neutral α-amylase, <u>A. niger</u> acid stable α-amylase, <u>A. niger</u> glucoamylase, <u>Rhizomucor miehei</u> lipase, <u>A. oryzae</u> alkaline protease or <u>A. oryzae</u> triose phosphate isomerase.

- 10 In particular when the host organism is <u>A. oryzae</u>, a preferred promoter for use in the process of the present invention is the <u>A. oryzae</u> TAKA amylase promoter as it exhibits a strong transcriptional activity in <u>A. oryzae</u>. The sequence of the TAKA amylase promoter appears from EP 238 023.
- 15 Termination and polyadenylation sequences may suitably be derived from the same sources as the promoter.

The techniques used to transform a fungal host cell may suitably be as described in EP 238 023.

To ensure secretion of the lipase A or the lipase variant from the host cell, the DNA sequence encoding the lipase variant may be preceded by a signal sequence which may be a naturally occurring signal sequence or a functional part thereof or a synthetic sequence providing secretion of the protein from the cell. In particular, the signal sequence may be derived from a 25 gene encoding an <u>Aspergillus</u> sp. amylase or glucoamylase, a gene encoding a <u>Rhizomucor miehei</u> lipase or protease, or a gene encoding a <u>Humicola</u> cellulase, xylanase or lipase. The signal sequence is preferably derived from the gene encoding <u>A. oryzae</u> TAKA amylase, <u>A. niger</u> neutral α-amylase, <u>A. niger</u> acid-stable 30 α-amylase or <u>A. niger</u> glucoamylase.

The medium used to culture the transformed host cells may be any conventional medium suitable for culturing <u>Aspergillus</u> cells. The transformants are usually stable and may be cultured in the absence of selection pressure. However, if the transformants are found to be unstable, a selection marker introduced into the cells may be used for selection.

The mature lipase protein secreted from the host cells may conveniently be recovered from the culture medium by well-known procedures including separating the cells from the medium by centrifugation or filtration, and precipitating proteinaceous components of the medium by means of a salt such as ammonium sulphate, followed by chromatographic procedures such as ion exchange chromatography, affinity chromatography, or the like.

It will be understood that the lipase variants of the invention 15 are contemplated to be active towards the same type of substrates as their parent lipases, with an improved specific activity. Thus, the lipase variants of the invention are contemplated to be useful for the same purposes as their parent lipases.

20 Accordingly, lipase variants of the invention prepared from a parent lipase useful as a detergent enzyme may be used as an active ingredient in a detergent additive or a detergent composition.

Another contemplated use of lipase variants of the invention, 25 is as digestive enzymes, e.g. in the treatment of cystic fibrosis.

A third use of the lipase variants of the invention, especially variants of <u>C. antarctica</u> lipases are in lipase-catalysed processes such as in ester hydrolysis, ester synthesis and interesterification. The use of lipases in these processes is discussed in detail in WO 88/02775 (Novo Nordisk A/S), the content of which is incorporated herein by reference. Further-

more, as the <u>C. antarctica</u> is an unspecific lipase, it may be used for randomization, e.g. in the preparation of margarine. Also the lipase variants of the invention may be used to avoid pitch trouble that arises in the production process for mechanical pulp or in a paper-making process using mechanical pulp, e.g. as described in PCT/DK92/00025 (Novo Nordisk A/S), the content of which is incorporated herein by reference.

BRIEF DESCRIPTION OF THE DRAWINGS

The present invention is described in the following with 10 reference to the appended drawings, in which

Fig. 1 is a computer model showing the three-dimensional structure of the lipid contact zone of the <u>H. lanuginosa</u> lipase described in WO 92/05249 when the lipase is in inactive (closed) and active (open) form, respectively. "White" residues 15 represent hydrophobic amino acids (Ala, Val, Leu, Ile, Pro, Phe, Trp, Gly and Met), "yellow" residues represent hydrophilic amino acids (Thr, Ser, Gln, Asn, Tyr and Cys), "blue" residues represent positively charged amino acids (Lys, Arg and His), and "red" residues represent negatively charged amino acids (Glu and Asp).

Figs. 2 and 3 illustrate the scheme for the construction of the expression plasmid pMT1229 (see Example 1).

The present invention is further illustrated in the following examples which are not intended, in any way, to limit the scope 25 of the invention as claimed.

MATERIALS

Plasmids and microorganisms

pBoel777 (p777) (described in EP 0 489 718)
p775 (the construction of which is described in EP 0 238 023)
5 pIC19H (Marsh et al., Gene 32 (1984), pp. 481-485)
pToC90 (described in WO 91/17243)
Aspergillus oryzae A1560: IFO 4177
E. coli MT172 (a K12 restriction deficient E. coli MC1000 derivative)

10 GENERAL METHODS

Site-directed in vitro mutagenesis of lipase genes

The three different approaches described in WO 92/05249 may be used for introducing mutations into the lipase genes, i.e. the oligonucleotide site-directed mutagenesis which is described by 20ller & Smith, DNA, Vol. 3, No. 6, 479-488 (1984), the PCR method as described in Nelson & Long, Analytical Biochemistry, 180, 147-151 (1989), and the so-called "cassette mutagenesis" technique, in which a segment between two restriction sites of the lipase-encoding region is replaced by a synthetic DNA fragment carrying the desired mutation. Use of the latter technique is illustrated in Example 2.

Determination of lipase specific activity

Lipase activity was assayed using glycerine tributyrat as a substrate and gum-arabic as an emulsifier. 1 LU (Lipase Unit) 25 is the amount of enzyme which liberates 1 μ mol titratable butyric acid per minute at 30°C, pH 7.0. The lipase activity was assayed by pH-stat using Radiometer titrator VIT90, Radiometer, Copenhagen. Further details of the assay are given in Novo Analytical Method AF 95/5, available on request.

EXAMPLES

EXAMPLE 1

Cloning of Candida antarctica lipase A

Chromosomal DNA of the <u>C. antarctica</u> strain LF058 (= DSM 3855 deposited with the Deutsche Sammlung von Mikroorganismen (DSM) on September 29, 1986 under the terms of the Budapest Treaty, and further described in WO 88/02775) was prepared by opening of frozen cells by grinding with quartz and subsequent extraction of DNA essentially as described by Yelton et al., (1984).

10 The purified DNA was cut partially with Sau3A and, after agarose gel electrophoresis, fragments in the range of 3-9 kb were isolated. The sized Sau3A fragments were ligated into a BamH1-cut, dephosphorylated plasmid pBR322 (New England Biolabs). The ligation mix was transformed into the <u>E. coli</u> MT172. Approximately 50,000 transformant <u>E. coli</u> colonies were obtained, 80% of which contained an insert of LF058 DNA.

Using standard colony hybridization techniques (Maniatis et al., 1982) the colonies were screened with the 32^P-phosphory-lated oligonucleotide probe NOR 440 (SEQ ID No. 7). NOR 440 is a degenerated (64) 17 mer based on the N-terminal determined from mature <u>C. antarctica</u> lipase A (SEQ ID No. 2). 34 colonies appeared positive after wash at low stringency (41°C and 6 x ssc). Plasmids were prepared from these colonies and Southern analyzed after restriction with BstN1. The probe for the southern was either the NOR 440 probe (SEQ ID No. 7) used for the colony hybridization (see above) or a 32^P-labelled probe NOR 438 (SEQ ID No. 6). NOR 438 is an oligonucleotide (a guess mer) where, at 13 positions, a base has been chosen on the basis of codon use in yeasts and filamentous fungi.

30 AACCCATACGACGACCC

T C T T T

NOR 440 (SEQ ID No. 7)

G T GCTGCTCTGCCTAACCCTTACGACGACCCTTTCTACACCACCCC NOR 438
T T T T (SEQ ID No. 6)

guess positions indicated

5 Only one plasmid, pMT1076, contained a band which hybridised both to NOR 440 at low stringency (see above) and to NOR 438 at a somewhat higher stringency (55°C and 1 x SSC).

PMT1076 was restriction mapped and the DNA sequence determined by the Maxam-Gilbert method. The sequence covering the open reading frame is shown in SEQ ID No. 1. The open reading frame is seen to encode a putative signal sequence of 21 amino acids (according to the von Heine rules (von Heijne, G. (1986)) and furthermore a propeptide of 10 amino acids preceding the N-terminal of the mature lipase. The last two amino acids of the propeptide are Arg Arg, i.e. a typical cleavage site for endoproteolytic processing by enzymes of the S. cereviciae KEX-2 type. The amino acid composition of the mature protein (starting at position 32) encoded by the DNA sequence is in agreement with the amino acid composition determined for C. antarctica lipase A, cf. the following table:

Amino acid composition of C. antartica lipase A (CALIP)

		Deduced from DNA sequence	By amino analysis	
5	Ala	50	47	
	Arg	9	9	
	Asp/AsN	35	36	
	Cys	4	. 4	
	Gln/GlN	35	36	
10	Gly	28	31	
	His	6	6	
	Ile	26	24	
	Leu	29	30	
	Lys	17	17	
15	Met	2 .	3	
	Phe	20	19	
	Pro	33	33	
	Ser	26	27	
	Thr	27	28	
20	Trp	5	4	
	Tyr	18	16	
	Val	27	26	

Through a number of standard plasmid manipulations (Maniatis et al., 1982) illustrated in Figs. 2 and 3, the open reading frame 25 of <u>C. antarctica</u> lipase A was placed in the correct orientation between the alpha-amylase promoter of <u>A. oryzae</u> and the glucoamylase transcription terminator of <u>A. niger</u>. The resulting expression plasmid pMT1229 was transformed into <u>A. oryzae</u> A1560 as described in EP 305,216. Transformants were isolated 30 and grown as described in the above cited patents and the culture supernatants were analyzed for the presence of <u>C. antarctica</u> lipase A.

EXAMPLE 2

Construction of a plasmid expressing the F135W variant of Candida_antarctica lipase A

A 246 bp BamHI/BssHII fragment was synthesized in vitro on the 5 basis of the nucleotide sequence of pMT1229 using oligonucleotide primers 3116 and 3117 in a PCR reaction. The primer 3117 includes a BssHII restriction site and a mutation in the 135 phe codon (TTC) to trp codon (TGG) which is marked with stars.

Oligonucleotide primer 3116 (F135W:256-276) (SEQ ID No. 8) 10 5'-CAG AAC GAG GCG GTG GCC GAC-3'

Oligonucleotide primer 3117 (F135W:566-487) (SEQ ID No. 9) 5'-TTC TTG AGC GCG CGG ATG CCG TCG AGG ATA GCC ATG CCC TCT TCG TAG CCA GCG ATG AAG GCG GCT TTC* C*AG CCT TCG TG-3'

The PCR reaction was performed by mixing the following com- 15 ponents and incubating the mixture in a HYBAIDTM thermal reactor.

	m		10 5-/47	1 μ1
	Template	PMT1229	10 ng/µl	
	H ₂ O			46.5µ1
	10 x PCR	buffer		10 <i>µ</i> l
20	2 mM dATE	•		10 <i>µ</i> l
	2 mM dTTF			10 μ1
	2 mM dCTI			10 µ1
	2 mM dGTI			10 µl
			50.5 pmol/	
	primer 31			
25	primer 31	L17	70.5 pmol/	
	Tag poly	nerase		0.5µl
	Parafin (50 μl
	Step I	94°C	2 min.	1 cycle
	Step II	• • •	30 sec.	-
30	_	50°C	30 sec.	30 cycle
30		72°C	2 min.	
				1 gyalo
	Step III	72°C	5 min.	1 cycle

The resulting 310 bp fragment was isolated from a 2% agarose gel after electrophoresis and digested with BamHI and BssHII 35 restriction enzymes. The resulting 264 bp BamHI/BssHII frag-

5

ment was likewise isolated from 2% agarose gel. This fragment was then ligated with

pMT1229	BamHI/XbaI	0.3	kb
pMT1229	BssHII/SphI	0.5	kb
pMT1229	SphI/XbaI	5.0	kb

The ligated DNA was transformed into E. coli strain MT172. Transformants which contained correct inserts were selected and their DNA sequence was determined by use of Sequenase (United States Biochemical Corporation). One resulting plasmid (pME-10 1178) contained a mutation in the amino acid position 135 (phe was mutated to trp).

pME1178 was cotransformed with pToC90 which included the amds gene from A. nidulans as a selective marker into the A. oryzae A1560 strain using the procedure described in WO 91/17243. A. 15 oryzae transformants were reisolated twice on selective plates and stable transformants were characterized by rocket immunoelectrophoresis, using anti-Candida lipase A antibody. Candida lipase A produced by a transformant (strain MEA65) was further analyzed for specific activity.

20 EXAMPLE 3

Construction of a plasmid expressing the F139W variant of Candida antarctica lipase A

A 246 bp BamHI/BssHII fragment was synthesized in vitro on the basis of the nucleotide sequence of the plasmid pMT1229 using 25 oligonucleotide primers 3116 and 3826 in a PCR reaction. The primer 3826 includes a BssHII restriction site and a mutation in the 139 phe codon (TTC) to trp codon (TGG) which is marked with stars.

Oligonucleotide primer 3116 is shown in Example 2.

Oligonucleotide primer 3826 (F139W:566-487) (SEQ ID No. 10) 5'-TTC TTG AGC GCG CGG ATG CCG TCG AGG ATA GCC ATG CCC TCT TCG TAG CCA GCG ATC* C*AG GCG GCT TTG AAG CCT TCG TG-3'

5 A PCR reaction was performed by the method described in Example 2. The 310 bp fragment was isolated from 2% agarose gel after electrophoresis and digested by BamHI and BssHII restriction enzymes. The resulting 264 bp BamHI/BssHII fragment was likewise isolated from 2% agarose gel. This fragment was then 10 ligated with

The ligated DNA was transformed into <u>E. coli</u> strain MT172.

15 Transformants which contained correct inserts were selected and their DNA sequence was determined by use of Sequenase (United States Biochemical Corporation). One resulting plasmid (pME-1229) contained a mutation in the amino acid position 139 (phe was mutated to trp).

20 pME1229 was cotransformed with pToC90 which included the amdS gene from A. nidulans as a selective marker into A. oryzae A 1560 strain. A oryzae transformants were reisolated twice on selective plates and enzyme activity of a stable transformant (MEA155) was analyzed by using tributylene as a substrate as 25 described in Example 5.

EXAMPLE 4

Construction of a plasmid expressing the F135W/F139W variant of Candida antarctica lipase A

A 246 bp BamHI/BssHII fragment was synthesized in vitro using 5 oligonucleotide primers 3116 and 4224 by a PCR reaction. The primer 4224 includes a BssHII restriction site and mutations in the 135 and 139 codons (TTC) to trp codons (TGG) which are marked with stars.

The oligonucleotide primer 3116 is shown in Example 2.

10 Oligonucleotide primer 4224 (F135W:566-487) (SEQ ID No. 11) 5'-TTC TTG AGC GCG CGG ATG CCG TCG AGG ATA GCC ATG CCC TCT TCG TAG CCA GCG ATC* C*AG GCG GCT TTC* C*AG CCT TCG TG-3'

PCR reaction was performed by using the method shown in Example 2. The 310 bp fragment was isolated from a 2% agarose gel after 15 electrophoresis and digested with BamHI and BssHII restriction enzymes. The resulting 264 bp BamHI/BssHII fragment was likewise isolated from a 2% agarose gel. This fragment was then ligated with

	pMT1229	BamHI/XbaI	0.3	kb
20	pMT1229	BssHII/SphI	0.5	kb
	pMT1229	SphI/XbaI	5.0	kb

The ligated DNA was transformed into E. coli MT172. Transformants which contained inserts were selected and their DNA sequence was determined by use of Sequenase. One resulting 25 plasmid (pME1230) contained two mutations in the amino acid positions 135 and 139 (phe was mutated to trp).

pME1230 was cotransformed with pToC90 which included the amdS gene from A. nidulans as a selective marker into A. oryzae A 1560 strain. A. oryzae transformants were reisolated twice on

selective plates and enzyme activity of stable transformants were analyzed by using tributylene as a substrate as described in Example 5.

EXAMPLE 5

5 Purification of <u>C. antarctica</u> lipase A variants F139W and F135W/F139W and comparison of specific activity with their parent wild type <u>C. antarctica</u> lipase A

The lipase variants and the parent lipase produced as described in Examples 3, 4 and 1, respectively, were purified using the 10 following 4 step standard purification procedure.

- Step 1: The fermentation broth containing the lipase and lipase variant, respectively, obtained by culturing the transformed A. oryzae cells described in Examples 1 and 3 above, was centrifuged, and the supernatant was adjusted to pH 7. Ionic strength was adjusted to 2 mSi. DEAE-Sephadex A-50 (Pharmacia) gel was swollen and equilibrated in 25 mM Tris acetate buffer pH 7. The fermentation supernatant was passed through DEAE-Sephadex A-50 on scintered glass funnel. The effluent containing lipase activity was collected and adjusted to 0.8 M ammonium acetate.
- 20 Step 2: An appropriate column was packed with TSK gel Butyl-Toyopearl 650 C and equilibrated with 0.8 M ammonium acetate. The effluent containing lipase activity was applied on the column. The bound material was eluted with water.
- Step 3: The lipase-containing eluate was then applied on a 25 Highperformance Q-Sepharose column. Lipase activity was collected as effluent. The lipases purified by this method were concentrated to an Optical Density of 1 at 280 nm.

The purity of the lipases was checked by SDS-PAGE showing one band with an molecular weight of about 45 kD. The lipase

activity was determined by use of the method outlined above in the section "General methods".

The lipase activity of the parent wild type lipase was 300 LU/OD₂₈₀ as compared to 1200 LU/OD₂₈₀ for the lipase variant 5 F139W. On the basis of OD₂₈₀ absorption without correction for the inserted tryptophan, the specific activity of the mutant was at least four times higher with the assay used. The lipase activity of the lipase variant F135W/F139W was 1400 LU/OD₂₈₀ (without correction for the two additional tryptophans).

10 EXAMPLE 6

Thermostability of Candida antarctica lipase A and the mutant F139W thereof

The thermostability of the C. antarctica lipase A and the C. antarctica lipase A variant, was examined by Differential 15 Scanning Calorimetry (DSC) at different pH values. Using this technique, the thermal denaturation temperature, T_d , is determined by heating an enzyme solution at a constant programmed rate.

More specifically, the Differential Scanning Calorimeter, MC20 2D, from MicroCal Inc. was used for the investigations. Enzyme solutions were prepared in 50 mM buffer solutions, cf. the tables below. The enzyme concentration ranged between 0.6 and 0.9 mg/ml, and a total volume of about 1.2 ml was used for each experiment. All samples were heated from 25°C to 90°C at a scan 25 rate of 90°C/hr.

The results obtained from the analysis is shown in the table below:

C. ant. lipase A (WT)

pН	Buffer (50 mM)	Denaturation temperature1)
4.5	Acetate	96°C
5 5	Acetate	95°C
7	TRIS	93°C

C. ant. lipase A mutant (F139W)

10 pH	Buffer (50 mM)	Denaturation temperature()
5	Acetate	84°C
7	TRIS	82°C

Temperature, at which approximately half the enzyme molecules present have been denatured thermally during heating

The above results show that the pH-optimum for the thermostability of *C. antarctica* lipase A and the F139W variant is unusually low and that both enzymes are very thermostable below pH 7. Within the investigated range the thermostability of both the Wild Type and the mutant F139W continues to increase as pH is lowered. This makes both lipases very well suited for hydrolysis/synthesis at unusually high temperatures at relatively low pH values.

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•	•	SEQUENCE LISTING	
	(1) GENER	RAL INFORMATION:	
5	(i)	APPLICANT: (A) NAME: NOVO NORDISK A/S (B) STREET: Novo Alle (C) CITY: Bagsvaerd (E) COUNTRY: DENMARK (F) POSTAL CODE (ZIP): DK-2880 (G) TELEPHONE: +45 44448888 (H) TELEFAX: +45 4449 3256 (I) TELEX: 37304	
	(ii)	TITLE OF INVENTION: Lipase Variants	
	(iii)	NUMBER OF SEQUENCES: 11	
15	(iv)	COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #	41 DI
20	(EPO)	(b) Software. Facencin Release #1.0, Version #	1.2
	(2) INFOR	RMATION FOR SEQ ID NO: 1:	
25	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 1389 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(iii)	HYPOTHETICAL: NO	
	(iii)	ANTI-SENSE: NO	
30	(v)	FRAGMENT TYPE: internal	
		ORIGINAL SOURCE: (A) ORGANISM: Candida antarctica (C) INDIVIDUAL ISOLATE: DSM 3855	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
35	ATGCGACTGT	OCTTGOGCTC CATCACGTCG CTGCTTGCGG CGGCAACGGC GGCTGTGCTC	60
	GOGGCTCCGG	OGGOOGAGAC GCTGGACOGA OGGGOGGOGC TGCCCAACCC CTACGACGAT	120

CCCTTCTACA CGACGCCATC CAACATCGGC ACGTTTGCCA AGGGCCAGGT GATCCAATCT

OSCAAGGIGC OCAOGGACAT OGGCAAGGCC AACAAGGCIG OGTOGITCCA GCIGCAGIAC

. •	OGCACCACCA	ATACGCAGAA	CEAGGCGGTG	GCCGACGTGG	CCACCGTGTG	GATCCCGGCC	300
	AAGCCCGCTT	CCCCCCCAA	GATCTTTTCG	TACCAGGICI	ACGAGGATGC	CACGGGGCTC	360
	GACTGTGCTC	CGAGCTACAG	CTACCTCACT	GGATTGGACC	AGCOGAACAA	GGTGACGGCG	420
	GTGCTCGACA	CCCCCATCAT	CATOGGCTGG	GOGCTGCAGC	AGGGCTACTA	OCTOCTOCTOC	480
5	TCCGACCACG	AAGGCTTCAA	AGCCGCCTTC	ATOGCTGGCT	ACGAAGAGGG	CATGGCTATC	540
	CTOGACGGCA	TCCGCGCGCT	CAAGAACTAC	CAGAACCIGC	CATCCGACAG	CAAGGTOGCT	600
	CTTGAGGGCT	ACACTGGCGG	AGCTCACGCC	ACCOTTCTGGG	CGACTTOGCT	TGCTGAATCG	660
	TACCCCCC	AGCTCAACAT	TGTCGGTGCT	TOGCACGGG	GCACGCCCCT	GAGCGCCAAG	720
	GACACCITTA	CATTCCTCAA	CGCCGACCC	Troscosser	TTGCCCTGGC	GCCICITICC	780
10	GCTCTCTCCC	TOGCTCATCC	TGATATGGAG	AGCTTCATTG	AGGCCCGATT	GAACGCCAAG	840
	GGTCAGCGGA	OCCICAAGCA	GATCCGCGC	CETCECTTCT	GCCTGCCGCA	CETEGRETTE	900
	ACCIACCCT	TCCTCAACCT	CITCICCIC	GTCAACGACA	CGAACCTGCT	GAATGAGGCG	960
	COGATOGCIA	GCATCCTCAA	GCAGGAGACT	GTGGTCCAGG	COGAAGOGAG	CTACACGGTA	1020
	TOGGTGCCCA	AGTTCCCGCG	CTTCATCTGG	CATGOGATOC	CCGACGAGAT	CETGCCGTAC	1080
15	CAGCCTGCGG	CTACCTACCT	CAAGGAGCAA	TGTGCCAAGG	GCGCCAACAT	CAATTITICG	1140
	CCCTACCCGA	TOGCOGAGCA	CCTCACCCC	GAGATCITTG	CICICGICCC	TAGCCTGTGG	1200
	TTTATCAAGC	AAGCCITCGA	OGGCACCACA	CCCAAGGTGA	TCIGCGGCAC	TCCCATCCCT	1260
	GCTATOGCTG	GCATCACCAC	GCCCTCGGCG	GACCAAGTGC	TGGGTTCGGA	CCTGGCCAAC	1320
	CAGCTGCGCA	GCCTCGACGG	CAAGCAGAGT	GOCTTOGGCA	AGCCCTTTGG	CCCCATCACA	1380
20	CCACCITAG						1389

(2) INFORMATION FOR SEQ ID NO: 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 463 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- 25

 - (ii) MOLECULE TYPE: protein
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
- 30 (v) FRAGMENT TYPE: N-terminal

(vi) ORIGINAL SOURCE: (A) ORGANISM: C

- (A) ORGANISM: Candida antarctica
- (C) INDIVIDUAL ISOLATE: DSM 3855
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
- 5 Met Arg Val Ser Leu Arg Ser Ile Thr Ser Leu Leu Ala Ala Ala Thr 1 5 10 15
 - Ala Ala Val Leu Ala Ala Pro Ala Ala Glu Thr Leu Asp Arg Ala 20 25 30

- Ala Leu Pro Asn Pro Tyr Asp Asp Pro Phe Tyr Thr Thr Pro Ser Asn 35 40 45
 - Ile Gly Thr Phe Ala Lys Gly Gln Val Ile Gln Ser Arg Lys Val Pro 50 60
 - Thr Asp Ile Gly Asn Ala Asn Asn Ala Ala Ser Phe Gln Leu Gln Tyr 65 70 75 80
- 15 Arg Thr Thr Asn Thr Gln Asn Glu Ala Val Ala Asp Val Ala Thr Val 85 90 95
 - Trp Ile Pro Ala Lys Pro Ala Ser Pro Pro Lys Ile Phe Ser Tyr Gln
 100 105 110
- Val Tyr Glu Asp Ala Thr Ala Leu Asp Cys Ala Pro Ser Tyr Ser Tyr 20 115 120 125
 - Leu Thr Gly Leu Asp Gln Pro Asn Lys Val Thr Ala Val Leu Asp Thr 130 135 140
 - Pro Ile Ile Ile Gly Trp Ala Leu Gln Gln Gly Tyr Tyr Val Val Ser 145 150 155 160
- 25 Ser Asp His Glu Gly Phe Lys Ala Ala Phe Ile Ala Gly Tyr Glu Glu 165 170 175
 - Gly Met Ala Ile Leu Asp Gly Ile Arg Ala Leu Lys Asn Tyr Gln Asn 180 185 190
- Leu Pro Ser Asp Ser Lys Val Ala Leu Glu Gly Tyr Ser Gly Gly Ala 30 195 200 205
 - His Ala Thr Val Trp Ala Thr Ser Leu Ala Glu Ser Tyr Ala Pro Glu 210 215 220
 - Leu Asn Ile Val Gly Ala Ser His Gly Gly Thr Pro Val Ser Ala Lys 225 230 235 240
- Asp Thr Phe Thr Phe Leu Asn Gly Gly Pro Phe Ala Gly Phe Ala Leu 245 250 255
 - Ala Gly Val Ser Gly Leu Ser Leu Ala His Pro Asp Met Glu Ser Phe

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		Ile	Glu	Ala 275	Arg	Leu	Asn	Ala	Lys 280	Gly	Gln	Arg	Thr	Leu 285	Lys	Gln	Ile
5		Arg	Gly 290	Arg	Gly	Phe	Cys	Leu 295	Pro	Gln	Val	Val	Leu 300	Thr	Tyr	Pro	Phe
		Leu 305	Asn	Val	Phe	Ser	Leu 310	Val	Asn	Asp	Thr	Asn 315	Leu	Leu	Asn	Glu	Ala 32 0
		Pro	Ile	Ala	Ser	Ile 325	Leu	Lys	Gln	Glu	Thr 330	Val	Val	Gln	Ala	Glu 335	Ala
ıo		Ser	Tyr	Thr	Val 340	Ser	Val	Pro	Lys	Phe 345	Pro	Arg	Phe	Ile	Trp 350	His	Ala
		Ile	Pro	Asp 355	Glu	Ile	Val	Pro	Tyr 360	Gln	Pro	Ala	Ala	Thr 365	Tyr	Val	Lys
15		Glu	Gln 370	_	Ala	Lys	Gly	Ala 375	Asn	Ile	Asn	Phe	Ser 380		Tyr	Pro	Ile
		Ala 385		His	Leu	Thr	Ala 390	Glu	Ile	Phe	Gly	Leu 395		Pro	Ser	Leu	Trp 400
-		Phe	Ile	Lys	Gln	Ala 405		Asp	Gly	Thr	Thr 410		Lys	Val	Ile	Cys 41 5	Gly
20		Thr	Pro	Ile	Pro 420		Ile	Ala	Gly	Ile 425		Thr	Pro	Ser	Ala 430		Gln
		Val	Leu	Gly 435		Asp	Leu	Ala	Asn 440		Leu	Arg	Ser	Leu 445		Gly	Lys
25		Glm	Ser 450		Phe	Gly	Lys	Pro 455		Gly	Pro	Ile	Thr 460		Pro	Glx	
	(2)	INI	FORM	ATI	ON E	OR	SEQ	ID	NO:	3:							
30		(:	i) s	(A) (B) (C) (D)	LEN TYI STI	IGTH PE: RAND	ARA : 2 ami EDN GY:	77 a no a ESS:	amin acid si	o a	cids	5					

- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO

- (v) FRAGMENT TYPE: internal 35
 - (vi) ORIGINAL SOURCE:

•		(A)	ORG	ANIS	M:	Pse	nobu	ona	s f	ragi	•				
	(xi) S	EQUE	NCE	DES	CRI	PTI	ON:	SEQ	ID	NO:	3:				
	Met Asp 1	Asp	Ser	Val 5	Asn	Thr	Arg	Tyr	Pro 10	Ile	Leu	Leu	Val	His 15	Gly
5	Leu Phe	Gly	Phe 20	Asp	Arg	Ile	Gly	Ser 25	His	His	Tyr	Phe	His 30	Gly	Ile
	Lys Gln	Ala 35	Leu	Asn	Glu	Cys	Gly 40	Ala	Ser	Val	Phe	Val 45	Pro	Ile	Ile
10	Ser Ala 50	Ala	Asn	Asp	Asn	Glu 55	Ala	Arg	Gly	Asp	Gln 60	Leu	Leu	Lys	Gln
	Ile His 65	Asn	Leu	Arg	Arg 70	Gln	Val	Gly	Ala	Gln 75	Arg	Val	Asn	Leu	Ile 80
	Gly His	Ser	Gln	Gly 85	Ala	Leu	Thr	Ala	Arg 90	Tyr	Val	Ala	Ala	Ile 95	Ala
15	Pro Glu	Leu	Ile 100	Ala	Ser	Val	Thr	Ser 105	Val	Ser	Gly	Pro	Asn 110	His	Gly
	Ser Glu	Leu 115	Ala	Asp	Arg	Leu	Arg 120	Leu	Ala	Phe	Val	Pro 125	Gly	Arg	Leu
20	Gly Glu 130		Val	Ala	Ala	Ala 135	Leu	Thr	Thr	Ser	Phe 140	Ser	Ala	Phe	Leu
	Ser Ala 145	Leu	Ser	Gly	His 150	Pro	Arg	Leu	Pro	Gln 155	Asn	Ala	Leu	Asn	Ala 160
	Leu Asn	Ala	Leu	Thr 165	Thr	Asp	Gly	Val	Ala 170	Ala	Phe	Asn	Arg	Gln 175	Tyr
25	Pro Gln	Gly	Leu 180	Pro	Asp	Arg	Trp	Gly 185	Gly	Met	Gly	Pro	Ala 190	Gln	Val
	Asn Ala	Val 195	His	Tyr	Tyr	Ser	Trp 200	Ser	Gly	Ile	Ile	Lys 205	Gly	Ser	Arg
30	Leu Ala 210		Ser	Leu	Asn	Leu 215		Asp	Pro	Leu	His 220	Asn	Ala	Leu	Arg
	Val Phe 225	Asp	Ser	Phe	Phe 230	Thr	Arg	Glu	Thr	Arg 235	Glu	Asn	Asp	Gly	Met 240
	Val Gly	Arg	Phe	Ser 245	Ser	His	Leu	Gly	Gln 250		Ile	Arg	Ser	Asp 255	Tyr
35	Pro Leu	Asp	His 260		Asp	Thr	Ile	Asn 265		Met	Ala	Arg	Gly 270		Ala

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Gly Ala Ser Thr Arg 275

- (2) INFORMATION FOR SEQ ID NO: 4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 269 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
- 10 (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (v) FRAGMENT TYPE: internal
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Rhizomucor miehei
- 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

Ser Ile Asp Gly Gly Ile Arg Ala Ala Thr Ser Gln Glu Ile Asn Glu 1 5 10 15

Leu Thr Tyr Tyr Thr Thr Leu Ser Ala Asn Ser Tyr Cys Arg Thr Val 20 25 30

20 Ile Pro Gly Ala Thr Trp Asp Cys Ile His Cys Asp Ala Thr Glu Asp 35 40 45

Leu Lys Ile Ile Lys Thr Trp Ser Thr Leu Ile Tyr Asp Thr Asn Ala 50 60

Met Val Ala Arg Gly Asp Ser Glu Lys Thr Ile Tyr Ile Val Phe Arg 25 65 70 75 80

> Gly Ser Ser Ser Ile Arg Asn Trp Ile Ala Asp Leu Thr Phe Val Pro 85 90 95

> Val Ser Tyr Pro Pro Val Ser Gly Thr Lys Val His Lys Gly Phe Leu 100 105 110

30 Asp Ser Tyr Gly Glu Val Gln Asn Glu Leu Val Ala Thr Val Leu Asp 115 120 125

> Gln Phe Lys Gln Tyr Pro Ser Tyr Lys Val Ala Val Thr Gly His Ser 130 135 140

Leu Gly Gly Ala Thr Ala Leu Leu Cys Ala Leu Gly Leu Tyr Gln Arg 35 145 150 155 160

Glu Glu Gly Leu Ser Ser Ser Asn Leu Phe Leu Tyr Thr Gln Gly Gln

		165		170	175
•	Dwo Aver Vol. Cla			Asn Tyr Val Val	
	180		185		190
5	Ile Pro Tyr Ard	g Arg Thr Va	al Asn Glu 200	Arg Asp Ile Val 205	
	Pro Pro Ala Ala 210		he Leu His 15	Ala Gly Glu Glu 220	Tyr Trp Ile
	Thr Asp Asn Sec 225	r Pro Glu T 230	hr Val Gln	Val Cys Thr Sea 235	Asp Leu Glu 240
10	Thr Ser Asp Cy	s Ser Asn S 245	er Ile Val	Pro Phe Thr Ser 250	r Val Leu Asp 255
	His Leu Ser Ty 26	_	le Asn Thr 265	Gly Leu Cys Se	£
(2)	INFORMATION	FOR SEQ I	D NO: 5:		
15	(i) SEQUENC	E CHARACT NGTH: 297			
	(B) TY	PE: amino RANDEDNES	acid		
		POLOGY: 1			
20	(ii) MOLECUI	E TYPE: p	rotein		
	(iii) HYPOTHE	TICAL: NO)		
	(iii) ANTI-SE	NSE: NO			
	(v) FRAGMEN	T TYPE: i	internal		
25	(- /	L SOURCE: RGANISM: P		niveus	
	(xi) SEQUENC	E DESCRIP	PTION: SE	Q ID NO: 5:	
	Asp Asp Asn La 1	u Val Gly (5	Gly Met Thr	Leu Asp Leu Pr 10	o Ser Asp Ala 15
30	Pro Pro Ile Se 20	_	Ser Ser Thr 25	Asn Ser Ala Se	er Asp Gly Gly 30
	Lys Val Val A 35	la Ala Thr !	Thr Ala Gln 40	n Ile Gln Glu Pr 45	_
	Ala Gly Ile A		Ala Tyr Cys 55	S Arg Ser Val Va 60	al Pro Gly Asn

Lys Trp Asp Cys Val Gln Cys Gln Lys Trp Val Pro Asp Gly Lys Ile 65 70 75 80

ė	Ile	Thr	Thr	Phe	Thr 85	Ser	Leu	Leu	Ser	Asp 90	Thr	Asn	Gly	Tyr	Val 95	Leu
	Arg	Ser	Asp	Lys 100	Gln	Lys	Thr	Ile	Tyr 105	Leu	Val	Phe	Arg	Gly 110	Thr	Asn
5	Ser	Phe	Arg 115	Ser	Ala	Ile	Thr	Asp 120	Ile	Val	Phe	Asn	Phe 125	Ser	Asp	Tyr
	Lys	Pro 130	Val	Lys	Gly	Ala	Lys 135	Val	His	Ala	Gly	Phe 140	Leu	Ser	Ser	Tyr
10	Glu 145	Gln	Val	Val	Asn	Asp 150	Tyr	Phe	Pro	Val	Val 155	Gln	Glu	Gln	Leu	Thr 160
	Ala	His	Pro	Thr	Tyr 165	Lys	Val	Ile	Val	Thr 170	Gly	His	Ser	Leu	Gly 175	Gly
	Ala	Gln	Ala	Leu 180	Leu	Ala	Gly	Met	Asp 185	Leu	Tyr	Gln	Arg	Glu 190		Arg
15	Leu	Ser	Pro 195		Asn	Leu	Ser	Ile 200		Thr	Val	Gly	Gly 205		Arg	Val
	Gly	Asn 210		Thr	Phe	Ala	Tyr 215		Val	Glu	Ser	Thr 220		Ile	Pro	Phe
20	Gln 225		Thr	Val	His	Lys 230		Asp	Ile	Val	Pro 235		Val	Pro	Pro	Gln 240
	Ser	Phe	Gly	Phe	Leu 245		Pro	Gly	Val	Glu 250		Trp	Ile	Lys	Ser 255	Gly
	Thr	Ser	Asn	Val 260		Ile	Cys	Thr	Ser 265		Ile	e Glu	Thr	Lys 270		Cys
25	Ser	: Asn	Ser 275		val	Pro	Ph∈	280		Ile	e Leu	a Asp	His 285		. Ser	Tyr
	Ph∈	ASP		Asr	Glu	Gly	Ser 295		Lev	ı						

(2) INFORMATION FOR SEQ ID NO: 6:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 44 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: DNA (synthetic) 35

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ŧ	(iii)	HYPOTHETICAL: NO	
	(iii)	ANTI-SENSE: NO	
	(v)	FRAGMENT TYPE: internal	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
5	GCIGCICICC	CIAACOCTIA OGAYGAYCCT TICIACACCA COCC	44
		RMATION FOR SEQ ID NO: 7:	
10	(1)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (synthetic)	
	(iii)	HYPOTHETICAL: NO	
	(iii)	ANTI-SENSE: NO	
15	(v)	FRAGMENT TYPE: internal	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
	AAYCCNTAYG	AYGAYCC	17
	(2) TNFO	RMATION FOR SEQ ID NO: 8:	
20	(1)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (synthetic)	
25	(iii)	HYPOTHETICAL: NO	
	(iii)	ANTI-SENSE: NO	
	(v)	FRAGMENT TYPE: internal	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 8:	
	CAGAACGAGG	COGTIGGOOGA C	21
30	(2) INFO	RMATION FOR SEQ ID NO: 9:	
	(i)	SEQUENCE CHARACTERISTICS:	

•		(A) LENGTH: 80 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
5	(ii)	MOLECULE TYPE: DNA (synthetic)	
	(iii)	HYPOTHETICAL: NO	
	(iii)	ANTI-SENSE: YES	
	(v)	FRAGMENT TYPE: internal	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 9:	
10	TTCTTGAGCG	CGCCGATGCC GTCGAGGATA GCCATGCCCT CTTCGTAGCC AGCGATGAAG	60
	GOGGCTTTCC	AGCCITOGIG	80
	(2) INFOR	RMATION FOR SEQ ID NO: 10:	
15	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 80 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (synthetic)	
	(iii)	HYPOTHETICAL: NO	
20	(iii)	ANTI-SENSE: YES	
	(v)	FRAGMENT TYPE: internal	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
	TTCTTGAGCG	OGOGGATGCC GTOGAGGATA GCCATGCCCT CTTOGTAGCC AGOGATCCAG	60
	GOGGCTTTGA	AGCCITCGIG	80
25	(2) INFO	RMATION FOR SEQ ID NO: 11:	
30		SEQUENCE CHARACTERISTICS: (A) LENGTH: 80 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (synthetic)	
	(iii)	HYPOTHETICAL: NO	

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į	(iii)	ANTI-SENSE: YES	
	(v)	FRAGMENT TYPE: internal	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
	TTCTTGAGCG	OGOGGATGCC GTOGAGGATA GOCATGCCCT CTTOGTAGCC AGOGATCCAG	60
5	GOGGCTTTCC	AGCCITCGIG	80

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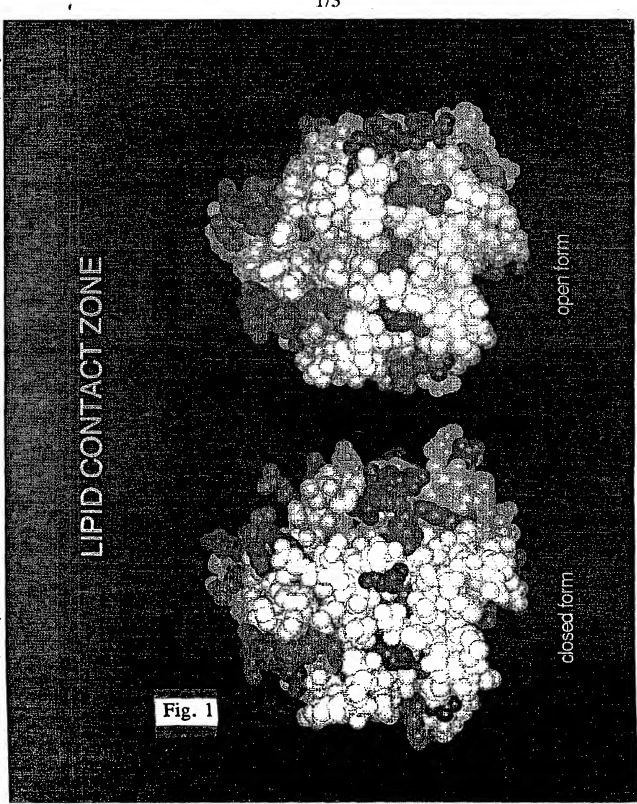
CLAIMS

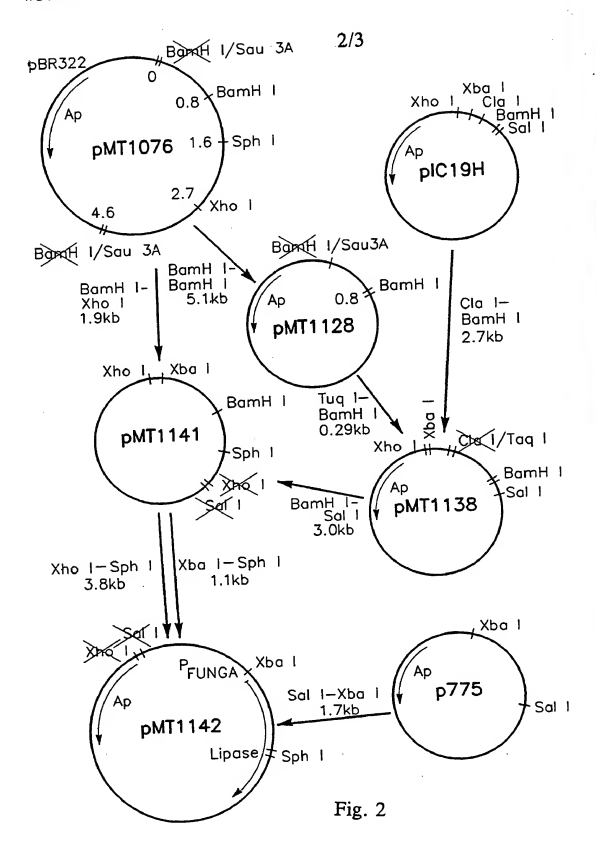
- 1. A lipase variant of a parent lipase comprising a trypsin-like catalytic triad including an active serine located in a predominantly hydrophobic, elongated binding pocket of the lipase molecule and, located in a critical position of a lipid contact zone of the lipase structure, an amino acid residue different from an aromatic amino acid residue, which amino acid residue interacts with a lipid substrate at or during hydrolysis, in which lipase variant said amino acid residue has been replaced by an aromatic amino acid residue so as to confer to the variant an increased specific activity as compared to that of the parent lipase.
- A lipase variant according to claim 1, wherein the aromatic amino acid residue to be inserted in the critical position is 15 selected from the group consisting of tryptophan, phenylalanine, tyrosine and histidine.
 - 3. A lipase variant according to claim 1 or 2, in which said amino acid residue different from an aromatic amino acid residue is a phenylalanine residue.
- 20 4. A lipase variant according to any of claims 1-3, in which the amino acid residue located in the critical position of the lipase is different from tryptophan, and said amino acid residue has been replaced with a tryptophan residue.
- A lipase variant according to any of the preceding claims,
 Wherein the parent lipase is selected from a microbial or a mammalian lipase.
 - 6. A lipase variant according to claim 5, wherein the parent lipase is a yeast lipase.
- 7. A lipase variant according to claim 6, wherein the parent 30 lipase is derived from a strain of <u>Candida antarctica</u>.

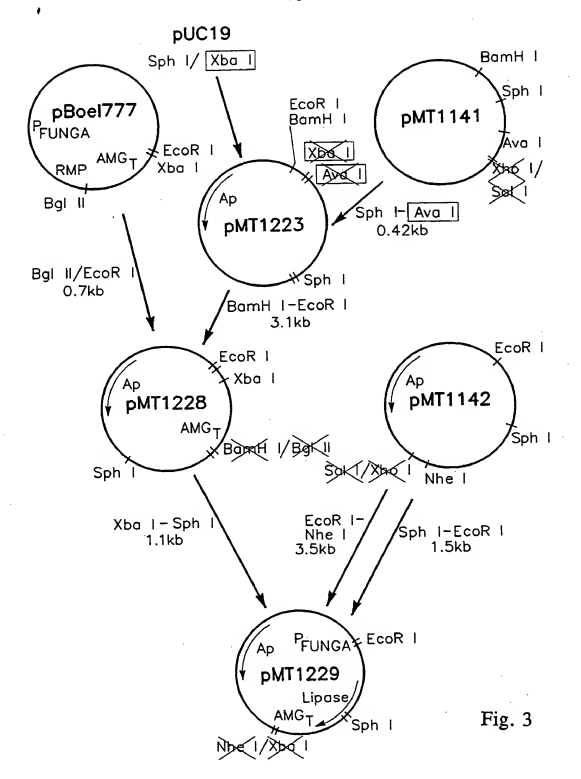
- , 8. A lipase variant according to claim 7, wherein the parent lipase is lipase A of <u>C. antarctica</u>.
- 9. A lipase variant according to claim 8, which has the amino acid sequence shown in SEQ ID No. 1, in which the phenylalanine 5 139 of the parent lipase has been replaced by a tryptophan residue, or in which the phenylalanine 135 and 139 of the parent lipase have been replaced by tryptophan residues.
 - 10. A lipase variant according to claim 5, in which the parent lipase is a bacterial lipase.
- 10 11. A lipase variant according to claim 10, wherein the parent lipase is derived from a strain of Pseudomonas.
 - 12. A lipase variant according to claim 11, which is derived from a strain of <u>Ps. fragi</u>.
- 13. A lipase variant according to claim 12, which has the amino 15 acid sequence shown in SEQ ID No. 3 in which the phenylalanine 29 of the parent lipase has been replaced by a tryptophan residue.
- 14. A lipase variant according to claim 5, wherein the parent lipase is selected from a fungal lipase, a human lipase, a 20 murine lipase, a rat lipase or a canine lipase.
 - 15. A <u>C. antarctica</u> lipase A essentially free from other substances from <u>C. antarctica</u>, which comprises the amino acid sequence shown in SEQ ID No. 2, or a variant of said lipase which
 - 25 1) has lipase activity,
 - 2) reacts with an antibody reactive with at least one epitope of <u>C. antarctica</u> lipase A having the amino acid sequence SEQ ID No. 2, and/or

- (3) is encoded by a nucleotide sequence which hybridizes with an oligonucleotide probe prepared on the basis of the full or partial nucleotide sequence shown in SEQ ID No. 1 encoding the C. antarctica lipase A.
- 5 16. A DNA sequence encoding <u>C. antarctica</u> lipase A having the amino acid sequence shown in SEQ ID No. 2 or a modification of said DNA sequence which encodes a variant of <u>C. antarctica</u> lipase A which
 - 1) has lipase activity,
- 10 2) reacts with an antibody reactive with at least one epitope of the <u>C. antarctica</u> lipase A having the amino acid sequence SEQ ID No. 2, and/or
- 3) is encoded by a nucleotide sequence which hybridizes with an oligonucleotide probe prepared on the basis of the full or 15 partial nucleotide sequence shown in SEQ ID No. 1 encoding the C. antarctica lipase A.
 - 17. A DNA construct comprising a DNA sequence encoding a lipase variant according to any of claims 1-14 or <u>C. antarctica</u> lipase A according to claim 15.
- 20 18. A recombinant expression vector which carries a DNA construct according to claim 17.
 - 19. A cell which is transformed with a DNA construct according to claim 17 or a vector according to claim 18.
- 20. A cell according to claim 19 which is a fungal cell, e.g. 25 belonging to the genus Aspergillus, such as A. niger, A. oryzae, or A. nidulans; a yeast cell, e.g. belonging to a strain of Saccharomyces, such as S. cerevisiae, or a methylotrophic yeast from the genera Hansenula, such as H. polymorpha, or Phichia, such as P. pastoris; or a bacterial cell,

- e.g. belonging to a strain of <u>Bacillus</u>, such as <u>B. subtilis</u>, <u>B. licheniformis</u> or <u>B. lentus</u>, or to a strain of <u>Escherichia</u>, such as <u>E. coli</u>.
- 21. A method of producing a lipase variant according to any of 5 claims 1-14, wherein a cell according to claim 19 or 20 is cultured under conditions conducive to the production of the lipase variant, and the lipase variant is subsequently recovered from the culture.
- 22. Use of a lipase variant according to any of claims 1-14 or 10 the <u>C. antarctica</u> lipase A or a variant thereof according to claim 15 in ester hydrolysis, ester synthesis or interesterification.
- 23. Use of a lipase variant according to any of claims 1-14 or the <u>C. antarctica</u> lipase A or a variant thereof according to 15 claim 15 for avoiding pitch trouble in a process for the production of mechanical pulp or a paper-making process using mechanical pulp.







International application No. PCT/DK 93/00225

A. CLASSIFICATION OF SUBJECT MATTER

IPC5: C12N 9/20, C12N 15/55 // (C 12 N 9/20, C 12 R 1:72)
According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC5: C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

SE,DK,FI,NO classes as above

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

WPI, CA, BIOSIS, MEDLINE

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	WO, A1, 9205249 (NOVO NORDISK A/S), 2 April 1992 (02.04.92), see claim 1	1-14,17-23
		
A	EP, A1, 0407225 (UNILEVER PLC ET AL), 9 January 1991 (09.01.91), see the claims	1-14,17-23
		
A	EP, A1, 0305216 (NOVO INDUSTRI A/S), 1 March 1989 (01.03.89)	1-14,17-23
		
Α .	NATURE, Volume 351, June 1991, Joseph D. Schrag et al, "Ser-His-Glu triad forms the catalytic site of the lipase from Geotrichum candidum" page 761 - page 764	1-14,17-23

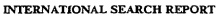
X Further documents are listed in the continuation of Box C. X See patent family annex.

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	•	Special	categories	of cited	documents

- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" ertier document but published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed
- T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- "X" document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
- "&" document member of the same patent family

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International application No. PCT/DK 93/00225

C (Continu	ation). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
A	NATURE, Volume 343, February 1990, Leo Brady et al, "A serine protease triad forms the catalytic centre of a triacylglycerol lipase" page 767 - page 770	1-14,17-23
		
Х	WO, A1, 8802775 (NOVO INDUSTRI A/S), 21 April 1988 (21.04.88), figure 1, claims 7,17, example 14-16	15-20-22-23
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x	Chemical Abstracts, Volume 118, No 9, 1 March 1993 (01.03.93), (Columbus, Ohio, USA), Patkar, S.A. et al, "Purification of two lipases from Candida antarctica and their inhibition by various inhibitors", page 321, THE ABSTRACT No 75881b, Indian J. Chem., Sect.B 1993, 32B (1), 76-80	15-20,22-23
		
A	Patent Abstracts of Japan, Vol 13,No 549, C-662, abstract of JP, A, 1-225481 (KURITA WATER IND LTD), 8 Sept 1989 (08.09.89)	15-20,22-23
A	Chemical Abstracts, Volume 106, No 21, 25 May 1987 (25.05.87), (Columbus, Ohio, USA), Omar, Ibrahim Che et al, "Purification and some properties of a thermostable lipase from Humicola lanuginosa No. 3", page 310, THE ABSTRACT No 171667j, Agric.Biol.Chem. 1987, 51 (1), 37-45	15-20,22-23
	<u></u>	
A	Chemical Abstracts, Volume 76, No 13, 27 March 1972 (27.03.72), (Columbus, Ohio, USA), Kosugi, Yoshiji et al, "Thermostable lipase form Pseudomonas species. Culture conditions and properties of the crude enzyme", page 267, THE ABSTRACT No 70997y, Hakko Kogaku Zasshi 1971, 49 (12), 968-980	15-20,22-23
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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

International application No. PCT/DK 93/00225

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
A		15-20,22-23
·	Chemical Abstracts, Volume 118, No 1, 4 January 1993 (04.01.93), (Columbus, Ohio, USA), Sugihara, Akio et al, "Purification and characterization of a novel ther mostable lipase from Pseudomonas cepacia", page 301, THE ABSTRACT No 2772g, J. Biochem. 1992, 112 (5),	
	598-603	
		
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International application No.

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Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
rnational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	
Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:	
Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	
ernational Searching Authority found multiple inventions in this international application, as follows:	١
aims 1-14,21 part of claims 17-20 and part of claims 22-23 directed a lipase variant	
aims 15-16, part of claims 17-20 and part of claims 22-23 directed to Candida antarctica lipase A	
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:	
	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). Observations where unity of invention is lacking (Continuation of item 2 of first sheet) creational Searching Authority found multiple inventions in this international application, as follows: Laims 1-14, 21 part of claims 17-20 and part of claims 22-23 directed a lipase variant Laims 15-16, part of claims 17-20 and part of claims 22-23 directed to Candida antarctica lipase A As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

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Information on patent family members

International application No.

PCT/DK 93/00225 16/10/93

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EP-A1-	0407225	09/01/91	JP-T- WO-A-	4500608 9100910	06/02/92 24/01/91	
EP-A1-	0305216	01/03/89	JP-A- JP-C- JP-B-	1157383 1761424 4038394	20/06/89 20/05/93 24/06/92	
WO-A1-	88027 7 5	21/04/88	EP-A- EP-A-	0287634 0382767	26/10/88 22/08/90	

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